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Compositions for diagnosis and therapy of diseases associated with aberrant expression of kremen and/or wnt

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Compositions for diagnosis and therapy of diseases associated with aberrant expression of kremen and/or wnt

The present invention relates to a composition useful for the diagnosis of diseases associated with aberrant expression of the gene encoding the receptor Kremen 1 and/or Kremen 2, e.g. tumors, diseases of the kidneys, bones and eyes. The present invention also relates to a pharmaceutical composition containing a compound which is capable of modifying (a) the expression of the gene encoding Kremen 1 and/or 2 or (b) the activity of the Kremen 1 and/or 2 receptor.

Tumorigenesis represents a complex multistage process in which genetic changes and environmental factors are thought to that control the cellular processes proliferation and differentiation. Among others the Wnt signal a crucial role as regards regulation cascade plays proliferation and differentiation of cells embryogenesis, as shown, e.g., in Drosophila, Xenopus and mice (Nusse and Varmus, Cell 69 (1992), 1073-1087). Wnt-genes encode secretory glycoproteins which activate a well characterized signal cascade via a Wnt receptor called "frizzled". The most prominent members of effectors of this signal cascade are betacatenin as wells the APC tumor suppressor gene (Miller and Moon, Genes Dev. 10 (1996), 2527-2539). Several indicate that an aberrant Wnt signal cascade might be involved in the development of colon cancer, breast cancer and melanoma (Pfeifer, Science, 275 (1997), 1752-1753; Polakis, Genes Dev. 14 (2000), 1837-1851). The first gene encoding a protein of the Wnt signal cascade, int-1, was isolated from mouse mammary tumor virus (MMTV) and it could be shown that it is an oncogene. It is assumed that an aberrant regulation of the activity of Wnt and/or components of the Wnt signal cascade downstream of the Wnt signal, e.g., beta-catenin and APC, are involved in tumorigenesis. In recent studies, a new family of genes, Dkk ("Dickkopf"), could be identified acting as

inhibitors of Wnt.

DKK1 binds and inhibits the Wnt coreceptor LRP 5/6 (Zorn, Curr. Biol. 11 (2001), R592-595) but otherwise, little is known about the mechanism of modulation of the Wnt signal cascade via Dkk. Accordingly, means for the therapy or diagnosis of diseases associated with a dis-regulated Wnt signal cascade were not Thus, the use of reliable diagnostic molecular markers would be helpful for an understanding of the molecular basis of diseases associated with an aberrant Wnt e.g., for distinguishing benign from cascade, e.g. tumors, malign tissue. In addition, Wnt signalling is involved in renal fibrosis (Surendran, Am J Physiol Renal Physiol 282 (2002) 431-441), polycystic kidney disease (Saadi-Kheddouci, Oncogene 20 (2001) 5972-5981) and the Dkk receptor LRP5 is involved in disorder osteoporosis-pseudoglioma recessive autosomal syndrome, whichh affects bones and eyes (OPPG; Gong, Cell (2001) $\underline{107}$, 513-523). It can be expected that such markers are also useful for therapy and for the development of novel therapeutic avenues for treatment of signal cascade Wnt dependent diseases, e.g. tumors or diseases of the kidneys, bones and eyes.

Thus, the technical problem underlying the present invention is to provide means for diagnosis and therapy of diseases associated with an aberrant Wnt signal cascade.

The solution to said technical problem is achieved by providing the embodiments characterized in the claims. During the experiments resulting in the present invention two genes, kremen 1 and 2, could be identified the products of which bind with high affinity to the polypeptides Dkkl and Dkk2. It could be shown that this binding is of physiological relevance since cotransfection of cells with dkkl as well as kremen 1 and 2 results in a synergistic inhibition of activation of the Wnt signal cascade. These data show that Kremen (1 and 2) can be

regarded as a receptor for the Dkk polypeptides and that the biological function of Kremen is the mediation of inhibition of the Wnt signal cascade via Dkk polypeptides. The data obtained provide evidence that the expression of kremen is very complex and that the genes encoding Kremen are involved in a variety of biological functions and might have tumor suppressor activity. Thus, Kremen is useful for the diagnosis and the development of therapies for Wnt mediated diseases. It can be expected that, e.g., the inhibition of the Wnt signal cascade by increasing the expression of kremen and/or by stimulating the activity of the polypeptide itself might have a therapeutic effect. On the other hand, the Kremen receptor (or the gene encoding it) can be regarded as a drug target allowing the identification of compounds useful for therapy.

BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1: Multisequence nucleic acid alignment of cDNAs encoding Kremenl (krm1) and 2 (krm2) from mouse and human hkrm1 and 2 are deduced from the human genome sequence in public data bases. Identical nucleotides are highlighted in black.

Figure 2: Multisequence amino acid alignment of Kremen1 and 2 proteins deduced from mouse and human cDNAs (see Figure 1)

Identical amino acids are highlighted in black, similar amino acids are in grey.

Figure 3: Kremen is a high affinity receptor for Dkkl and Dkk2 293T cells were transfected with cytomegalovirus (CMV) promoter-driven expression plasmids encoding mkrml (top) or mkrm2 (bottom) as indicated, incubated with recombinant Dkkl-AP, Dkk2-AP or Dkk3-AP and stained for bound AP activity. TOP: Binding curves and Scatchard analysis of Dkk-AP fusion proteins

binding to mkrm2 transfected cells. Bottom: Binding curves for Dkk-APs binding to mkrm1 transfected cells. Dissociation constants (K_d) are indicated; a, c: Binding curves; b, d, e: Scatchard analysis.

Figure 4: Kremen and Dkkl synergistically inhibit the Wnt signal cascade

293 kidney cells were transfected with the Wnt reporter (TOP-FLASH) with or without the genes indicated. Two days after transfection, the luciferase activity expressed was determined. RLU: relative light units (normalized against cotransfected Renilla luciferase). Xdkkl = Xenopus dkkl; mkrml; 2 = mouse kremen 1,2; wnt = mouse wntl; fz = mouse frizzled8; lrp6 = human lrp6.

Figure 5: Expression of kremen in mice

The expression of kremen 1 and kremen 2 was analysed by RT-PCR in various tissues of adult mice. The results were normalized using constitutive histon H4 expression. Abbreviations: -RT= control sample in which reverse transcriptase was omitted; sk muscle= skelletal muscle; mam. gland= mammary gland; H4= Histone 4 as loading control; mkrm1, 2= mouse kremen 1, 2.

The present invention relates to a diagnostic composition comprising (a) at least one nucleic acid molecule which is capable of specifically hybridizing to the nucleotide sequence encoding Kremen 1 as depicted in Figure 1 and/or to the nucleotide sequence encoding Kremen 2 as depicted in Figure 2, or (b) at least one ligand which is capable of specifically binding to a Kremen 1 and/or Kremen 2 polypeptide.

As used herein the term "Kremen 1 polypeptide" and "Kremen 2 polypeptide" not only refers to polypeptids encoded by the nucleotide sequence as depicted in Figure 1 and/or 2 but also to

polypeptides differing in amino acid sequence due to insertion, deletion and/or substitution of one ore more amino acids and showing at least one biological activity of a Kremen 1 and/or Kremen 2 receptor, e.g. the ability of signal transduction after ligand binding. Preferably, the related polypeptides are polypeptides the amino acid sequence of which shows an identity of at least 40%, in particular an identity of at least 65%, preferably of at least 80% and, particularly preferred, of at least 90% to the amino acid sequences of the polypeptides encoded by the nucleotide sequences shown in Figure 1 or 2.

The nucleic acid molecules useful as probes can be both DNA and RNA molecules, preferably they are single-stranded DNA molecules. They can be isolated from natural sources or can be synthesized according to know methods.

As a hybridization probe nucleic acid molecules can be used, for example, that have a nucleotide sequence which is exactly or basically complementary to a nucleotide sequence as depicted in Figure 1 and 2, respectively, or parts of these sequences. The fragments used as hybridization probe can be synthetic fragments that were produced by means of conventional synthetic methods

As used herein, the term "hybridizing, relates to hybridization under conventional hybridization conditions, preferably under stringent conditions as described, for example, in Sambrook et al., Molecular Cloning, A Laboratory Manual 2nd edition (1989) Cold Spring Harbor Laboratory Press, Cold Spring Harbor, NY. However, in certain cases, a hybridizing nucleic acid molecule can also be detected at lower stringency hybridization conditions. Changes in the stringency of hybridization and signal detection are primarily accomplished through the manipulation of formamide concentration (lower percentages of formamide result in lowered stringency), salt conditions, or temperature. For example, lower stringency conditions include

an overnight incubation at 37°Cin a solution comprising 6X SSPE (20X SSPE = 3M NaCl; 9.2M NaH₂PO₄; 0.02M EDTA, pH7.4), 0.5% 30% formamide, 100 µg/ml salmon sperm blocking DNA, following by washes at 50°C with 1 X SSPE, 0.1% SDS. addition, to achieve even lower stringency, washes performed following stringent hybridization can be done at higher salt the above Variations in SSC). 5X concentrations (e.g. conditions may be accomplished through the inclusion and/or substitution of alternate blocking reagents used to suppress background in hybridization experiments. The inclusion of specific blocking reagents may require modification of the hybridization conditions described above, due to problems with compatibility.

The term "ligand" as used herein refers to any molecule which is capable of specifically binding to Kremen 1 and/or Kremen 2, thus allowing to determine the level of receptor molecules. antibodies, include molecules such Examples oligonucleotides, proteins or small molecules. The molecule can be the natural ligand of Kremen, i.e. Dkk1 or Dkk2, or can be closely related to said ligand, e.g., a fragment of the ligand, or a natural substrate, a ligand, a structural or functional mimetic; see, e.g., Coligan, Current Protocols in Immunology 1(2) (1991); Chapter 5. In either case, the molecule can be isolated or rationally designed using known techniques; see also infra.

Preferably, the ligand is an antibody. The term "antibody", preferably, relates to antibodies which consist essentially of different epitopic with antibodies monoclonal pooled antibody. monoclonal well distinct as specifities, as preparations. Monoclonal antibodies are made from an antigen containing fragments of Kremen 1 or Kremen 2 by methods well known to those skilled in the art (see, e.g., Köhler et al., Nature 256 (1975), 495). As used herein, the term "antibody" (Ab) or "monoclonal antibody" (Mab) is meant to include intact

molecules as well as antibody fragments (such as, for example, Fab and F(ab') 2 fragments) which are capable of specifically binding to Kremen. Fab and f(ab')2 fragments lack the Fc fragment of intact antibody, clear more rapidly from the circulation, and may have less non-specific tissue binding than an intact antibody. (Wahl et al., J. Nucl. Med. 24: 316-325 (1983)). Thus, these fragments are preferred, as well as the products of a FAB or other immunoglobulin expression library. Moreover, antibodies of the present invention include chimerical, single chain, and humanized antibodies.

For certain purposes, e.g. diagnostic methods, the nucleic acid molecule used as probe or the ligand, e.g., antibody, can be detectably labeled, for example, with a radioisotope, a bioluminescent compound, a chemiluminescent compound, a fluorescent compound, a metal chelate, or an enzyme.

The nucleic acid molecules can be used, for example, as probes or primers in the diagnostic assays described below and allow, e.g., the analysis of the expression of kremen 1 and 2 by determining the mRNA level or the determination of mutations within the coding region or regulatory regions leading to polypeptide molecules with altered, e.g. destroyed, activity, or leading to altered expression. Preferably, the nucleic acid molecules are oligonucleotides having a length of at least 10, in particular of at least 15 and particularly preferred of at least 50 nucleotides. These nucleic acid molecules of the invention can also be used, for example, as primers for a PCR reaction.

The present invention also relates to the use of a nucleic acid molecule or ligand as defined above for the preparation of a diagnostic composition for the diagnosis of a disease associated with (a) aberrant expression of kremen 1 and/or kremen 2 and/or (b) aberrant activity of a Kremen 1 and/or Kremen 2 polypeptide.

In a preferred embodiment, the target to which the nucleic acid molecule hybridizes is an mRNA.

The present invention also provides a method of diagnosing a disease associated with (a) aberrant expression of kremen 1 and/or kremen 2 and/or (b) aberrant activities or amounts of a Kremen 1 and/or Kremen 2 polypeptide in a subject comprising:

- (a) determining (a) the amount of expression of kremen 1 and/or kremen 2 and/or (b) the amount of biologically active Kremen 1 and/or Kremen 2 polypeptide in a biological sample; and
- (b) diagnosing a disease associated with (a) aberrant expression of kremen 1 and/or kremen 2 and/or (b) aberrant activites or amounts of a Kremen 1 and/or Kremen 2 polypeptide or a risk for the development of such diasease based on an altered amount of expression of kremen 1 and/or kremen 2 and/or (b) altered activities or amounts of biologically active Kremen 1 and/or Kremen 2 polypeptide compared to a control.

Suitable assay formats are well known to the person skilled in the art and, in addition, described below. Suitable positive control samples expressing human kremen 1 and 2 protein are, e.g., HEK 293 cells.

The Kremen 1 or 2 polypeptide or the corresponding mRNA, e.g. in biological fluids or tissues, may be detected directly in situ, e.g. by in situ hybridization or it may be isolated from other cell components by common methods known to those skilled in the art before contacting with a probe. Detection methods include Northern Blot analysis, RNase protection, methods, e.g. in situ hybridization, in vitro amplification methods (PCR, LCR. QRNA replicase or RNAtranscription/amplification (TAS, 3SR), reverse dot blot disclosed in EP-B1 O 237 362), immunoassays, Western Blot and

other detection assays that are known to those skilled in the art.

specific specific antibody or (e.g. а The diagnostic composition can of the oligonucleotide) detectably labeled. In a preferred embodiment, said diagnostic composition contains an anti-Kremen 1 or -Krmen-2 antibody and allows said diagnosis, e.g., by ELISA and contains the antibody bound to a solid support, for example, a polystyrene microtiter dish or nitrocellulose paper, using techniques known in the art. Alternatively, said diagnostic compositions are based on a RIA and contain said antibody marked with a radioactive isotope. Suitable antibody assay labels are known in the art and include enzyme labels, such as, glucose oxidase, radioisotopes, such as iodine (125I, 121I), carbon (14C), sulfur (^{35}S) , tritium (^{3}H) , indium (^{112}In) , and technetium rhodamine, and biotin. In addition to assaying Kremen levels biological sample, the polypeptide can also be detected in vivo by imaging. Antibody labels or markers for in vivo imaging of protein include those detectable by X-radiography, NMR or ESR. For X-radiography, suitable labels include radioisotopes such as barium or cesium, which emit detectable radiation but are not overtly harmful to the subject. Suitable markers for NMR and ESR include those with a detectable characteristic spin, such as deuterium, which may be incorporated into the antibody by labeling of nutrients for the relevant hybridoma. A proteinspecific antibody or antibody fragment which has been labeled with an appropriate detectable imaging moiety, such as a radioisotope (for example, 131 I, 112 In, 99 mTc), a radio-opaque by nuclear a material detectable substance, or parenterally, example, introduced (for is resonance, subcutaneously, or intraperitoneally) into the mammal. It will be understood in the art that the size of the subject and the imaging system used will determine the quantity of imaging moiety needed to produce diagnostic images. In the case of a radioisotope moiety, for a human subject, the quantity of radioactivity injected will normally range from about 5 to 20 millicuries of ⁹⁹mTc. The labeled antibody or antibody fragment will then preferentially accumulate at the location of cells which contain the specific Kremen polypeptide. In vivo tumor imaging is, e.g., described in S.W. Burchiel et al., "Immunopharmacokinetics of Radiolabeled Antibodies and Their Fragments,". (Chapter 13 in Tumor Imaging: The Radiochemical Detection of Cancer, S.W. Burchiel and B.A. Rhodes, eds., Masson Publishing Inc. (1982)).

In a further aspect, the present invention, relates to a method for identifying a binding partner to a Kremen 1 and/or 2 polypeptide comprising:

- (a) contacting said polypeptide with a compound to be screened; and
- (b) determining whether the compound effects an activity of the polypeptide.

The invention also includes a method of identifying compounds which bind to a Kremen 1 and/or Kremen 2 polypeptide comprising the steps of:

- (a) incubating a candidate binding compound with said polypeptide; and
- (b) determining if binding has occurred.

Kremen 1 or 2 polypeptides may be used to screen for proteins or other compounds that bind to Kremen 1 or 2 or for proteins or other compounds to which Kremen 1 and 2 bind. The binding of Kremen 1 or 2 and the molecule may activate (agonist), increase, inhibit (antagonist), or decrease activity of Kremen 1 or Kremen 2 or the molecule bound. Examples of such molecules include antibodies, oligonucleotides, proteins (e.g., ligands), or small molecules.

Preferably, the molecule is closely related to the natural ligand of Kremen 1 or 2, e.g., a fragment of the ligand, or a

natural substrate, a ligand, a structural or functional mimetic; see, e.g., Coligan, Current Protocols in Immunology 1(2) (1991); Chapter 5.

Preferably, the screening for these molecules involves producing appropriate cells which express Kremen 1 and/or, either as a secreted protein or on the cell membrane. Preferred cells include cells from mammals, yeast, Drosophila, or E. coli. Cells expressing Kremen 1 and/or 2 (or cell membrane containing the expressed polypeptide) are then preferably contacted with a test compound potentially containing the molecule to observe binding, stimulation, or inhibition of activity of Kremen 1 and/or 2.

The assay may simply test binding of a candidate compound to Kremen 1 and/or 2, wherein binding is detected by a label, or in an assay involving competition with a labeled competitor. Further, the assay may test whether the candidate compound results in a signal generated by binding to Kremen 1 and/or Kremen 2. Suitable assays to analyze the activity of kremen 1 and /or 2 include Wnt-inducible luciferase reporter assays in transfected HEK 293 cells, where dkkl synergizes with kremen 1 and /or 2 to inhibit a Wnt1-induced signal, such as is shown in Figure 4.

Alternatively, the assay can be carried out using cell-free preparations, polypeptide/molecule affixed to a solid support, chemical libraries, or natural product mixtures. The assay may also simply comprise the steps of mixing a candidate compound with a solution containing Kremen 1 and/or Kremen 2, measuring Kremen/molecule activity or binding, and comparing the Kremen/molecule activity or binding to a standard.

Preferably, an ELISA assay can measure Kremen 1 and/or Kremen 2 level or activity in a sample (e.g., biological sample) using a monoclonal or polyclonal antibody. The antibody can measure

Kremen 1 and/or Kremen 2 level or activity by either binding, directly or indirectly, to Kremen 1 and/or Kremen 2 or by competing with Kremen 1 and/or Kremen 2 for a substrate. All of these above assays can be used as diagnostic or prognostic markers. The molecules discovered using these assays can be used to treat disease or to bring about a particular result in elimination (e.g., of а tumor, support patient etc.) by modulating, preferably regenerative processes activating the Kremen 1 and/or Kremen 2 molecule. Moreover, the assays can discover agents which may inhibit or enhance the 1 and/or Kremen 2 from suitably production of Kremen manipulated cells or tissues.

Moreover, the invention includes a method of identifying activators/agonists or inhibitors/antagonists of a Kremen 1 and/or Kremen 2 polypeptide comprising the steps of:

- (a) incubating a candidate compound with said polypeptide;
- (b) assaying a biological activity, and
- (c) determining if a biological activity of said polypeptide has been altered.

Suitable assays include analysis of formation of a ternary complex between kremen1 or kremen 2 with recombinant Dkk1 protein and recombinant extracellular domain of LRP6.

In a further embodiment, the present invention relates to method of identifying and obtaining a drug candidate for therapy of diseaseas associated with (a) aberrant expression of kremen 1 and/or kremen 2 and/or (b) aberrant activities or amounts of a Kremen 1 and/or Kremen 2 polypeptide comprising the steps of

(a) contacting a Kremen 1 and/or Kremen 2 polypeptide or a cell expressing said polypeptide, and optionally the corresponding ligand(s), in the presence of components capable of providing a detectable signal in response to binding to said drug candidate to be screened; and

(b) detecting presence or absence of a signal or increase of the signal generated, wherein the presence or increase of the signal is indicative for a putative drug.

Suitable assays to analyze the activity of kremen 1 and /or 2 include Wnt-inducible luciferase reporter assays in transfected HEK 293 cells, where dkkl synergizes with kremen 1 and /or 2 to inhibit a Wntl-induced signal, such as is shown in Figure 4.

The drug candidate may be a single compound or a plurality of compounds. The term "plurality of compounds" in a method of the invention is to be understood as a plurality of substances which may or may not be identical.

Said compound or plurality of compounds may be chemically synthesized or microbiologically produced and/or comprised in, for example, samples, e.g., cell extracts from, e.g., plants, animals or microorganisms. Furthermore, said compound(s) may be known in the art but hitherto not known to be capable of Kremen 1 and/or activating Kremen suppressing or polypeptides. The reaction mixture may be a cell free extract or may comprise a cell or tissue culture. Suitable set ups for the method of the invention are known to the person skilled in the art and are, for example, generally described in Alberts et al., Molecular Biology of the Cell, third edition (1994) and in the appended examples. The plurality of compounds may be, e.g., added to the reaction mixture, culture medium, injected into a cell or otherwise applied to a transgenic animal. The cell or tissue that may be employed in the method of the invention preferably is a host cell, mammalian cell or non-human transgenic animal.

If a sample containing a compound or a plurality of compounds is identified in the method of the invention, then it is either possible to isolate the compound from the original sample identified as containing the compound capable of suppressing or

activating a Kremen 1 and/or Kremen 2 polypeptide, or one can further subdivide the original sample, for example, consists of a plurality of different compounds, so as to reduce the number of different substances per sample and repeat the method with the subdivisions of the original sample. Depending on the complexity of the samples, the steps described above can several times, preferably until the performed identified according to the method of the invention only limited number of or only one substance(s). comprises a Preferably said sample comprises substances of similar chemical and/or physical properties, and most preferably said substances are identical.

Several methods are known to the person skilled in the art for producing and screening large libraries to identify compounds. having specific affinity for a target. These methods include the phage-display method in which randomized peptides displayed from phage and screened by affinity chromatography to an immobilized receptor; see, e.g., WO 91/17271, WO 92/01047, US-A-5,223,409. In another approach, combinatorial libraries of synthesized immobilized on а chip are polymers photolithography; see, e.g., US-A-5,143,854, WO 90/15070 and WO 92/10092. The immobilized polymers are contacted with a labeled receptor and scanned for label to identify polymers binding to the receptor. The synthesis and screening of peptide libraries on continuous cellulose membrane supports that can be used for identifying binding ligands of the Kremen polypeptides and, thus, possible inhibitors and activators is for example, in Kramer, Methods Mol. described, (1998), 25-39. This method can also be used, for example, for determining the binding sites and the recognition motifs in the Kremen 1 and/or 2 polypeptide. In like manner, the substrate specificity of the DnaK chaperon was determined and the contact interleukin-6 and its receptor; sites between human Rudiger, EMBO J. 16 (1997), 1501-1507 and Weiergraber, FEBS (1996), 122-126, respectively. Furthermore, 379

above-mentioned methods can be used for the construction of binding supertopes derived from the Kremen 1 or Kremen 2 polypeptide. A similar approach was successfully described for peptide antigens of the anti-p24 (HIV-1) monoclonal antibody; see Kramer, Cell 91 (1997), 799-809. A general route to fingerprint analyses of peptide-antibody interactions using the clustered amino acid peptide library was described in Kramer, Mol. Immunol. 32 (1995), 459-465. In addition, antagonists of a Kremen 1 and/or Kremen 2 polypeptide can be derived and identified from monoclonal antibodies that specifically react with a Kremen 1 and/or Kremen 2 polypeptide in accordance with the methods as described in Doring, Mol. Immunol. 31 (1994), 1059-1067.

All these methods can be used in accordance with the present invention to identify activators/agonists and inhibitors/antagonists of a Kremen 1 and/or Kremen 2 polypeptide.

Various sources for the basic structure of such an activator or inhibitor can be employed and comprise, for example, mimetic analogs of a Kremen 1 and/or Kremen 2 polypeptide. Mimetic analogs of a Kremen 1 and/or Kremen 2 polypeptide or biologically active fragments thereof can be generated by, for example, substituting the amino acids that are expected to be essential for the biological activity with, e.g., stereoisomers, i.e. Damino acids; see e.g., Tsukida, J. Med. Chem. 40 (1997), 3534-3541. Furthermore, in case fragments are used for the design of active analogs pro-mimetic components can biologically incorporated into a peptide to reestablish at least some of the conformational properties that may have been lost upon removal of part of the original polypeptide; see, e.g., Nachman, Regul. Pept. 57 (1995), 359-370. Furthermore, a Kremen 1 and/or Kremen 2 polypeptide can be used to identify synthetic chemical peptide mimetics that bind to or can function as a ligand, substrate or binding partner of said polypeptide(s) as effectively as does the natural polypeptide; see, e.g., Engleman, J. Clin. Invest. (1997), 2284-2292. For example, folding simulations and computer redesign of structural motifs of a Kremen 1 and/or Kremen 2 polypeptide can be performed using appropriate computer programs (Olszewski, Proteins 25 (1996), 286-299; Hoffman, Comput. Appl. Biosci. 11 (1995), 675-679). Computer modeling of protein folding can be used for the conformational and energetic analysis of detailed peptide and protein models (Monge, J. Mol. Biol. (1995), 995-1012; Renouf, Adv. Exp. Med. Biol. 376 (1995), 45). In particular, the appropriate programs can be used for the identification of interactive sites of a Kremen 1 and/or Kremen 2 polypeptide and its ligand or other interacting proteins by computer assistant searches for complementary peptide sequences Immunomethods 5 (1994), 114-120. Further appropriate (Fassina, computer systems for the design of protein and peptides are described in the prior art, for example in Berry, Biochem. Soc. Trans. 22 (1994), 1033-1036; Wodak, Ann. N. Y. Acad. Sci. Biochemistry 25 (1986), 5987-5991. The 1-13; Pabo, (1987), results obtained from the above-described computer analysis can be used for, e.g., the preparation of peptide mimetics of a Kremen 1 and/or Kremen 2 polypeptide or fragments thereof. Such pseudopeptide analogues of the natural amino acid sequence of the protein may very efficiently mimic the parent protein (Benkirane, example, (1996), 33218-33224). For Biol. Chem. 271 J. incorporation of easily available achiral ω -amino acid residues into a Kremen 1 or 2 polypeptide or a fragment thereof results in the substitution of amide bonds by polymethylene units of an aliphatic chain, thereby providing a convenient strategy for constructing a peptide mimetic (Banerjee, Biopolymers 39 (1996), 769-777). Superactive peptidomimetic analogues of small peptide hormones in other systems are described in the prior art (Zhang, Biochem. Biophys. Res. Commun. 224 (1996), 327-331). Appropriate peptide mimetics of a Kremen 1 and/or Kremen 2 polypeptide can peptide mimetic identified by the synthesis of be combinatorial libraries through successive amide alkylation and testing the resulting compounds, e.g., for their binding and immunological properties. Methods for the generation and use of peptidomimetic combinatorial libraries are described in the prior art, for example in Ostresh, Methods in Enzymology 267 (1996), 220-234 and Dorner, Bioorg. Med. Chem. 4 (1996), 709-715. Furthermore, a three-dimensional and/or crystallographic structure of a Kremen 1 and/or Kremen 2 polypeptide can be used for the design of peptide mimetic inhibitors of the biological activity of the polypeptide (Rose, Biochemistry 35 (1996), 12933-12944; Rutenber, Bioorg. Med. Chem. 4 (1996), 1545-1558).

It is also well known to the person skilled in the art, that it is possible to design, synthesize and evaluate mimetics of small organic compounds that, for example, can act as a substrate or ligand to a Kremen 1 and/or Kremen 2 polypeptide. For example, it has been described that D-glucose mimetics of hapalosin exhibited similar efficiency as hapalosin in antagonizing multidrug resistance assistance-associated protein in cytotoxicity; see Dinh, J. Med. Chem. 41 (1998), 981-987.

The nucleic acid molecule encoding a Kremen 1 and/or Kremen 2 polypeptide can also serve as a target for activators and inhibitors. Activators may comprise, for example, proteins that bind to the mRNA of a gene encoding a Kremen 1 and/or Kremen 2 polypeptide, thereby stabilizing the native conformation of the mRNA and facilitating transcription and/or translation, e.g., in like manner as Tat protein acts on HIV-RNA. Furthermore, methods are described in the literature for identifying nucleic acid molecules such as an RNA fragment that mimics the structure of a defined or undefined target RNA molecule to compound binds inside of a cell resulting in retardation of cell growth or cell death; see, WO 98/18947 and references cited therein. These nucleic acid molecules can be used for identifying unknown compounds of RNA interest, and for identifying unknown pharmaceutical targets for use in treating a disease. These methods and compositions can be used in screening for novel or for

identifying compounds useful to alter expression levels of polypeptids encoded by a nucleic acid molecule. Alternatively, for example, the conformational structure of the RNA fragment which mimics the binding site can be employed in rational drug design to modify known drugs to make them bind more avidly to the target. One such methodology is nuclear magnetic resonance (NMR), which is useful to identify drug and RNA conformational structures. Still other methods are, for example, the drug design methods as described in WO 95/35367, US-A-5,322,933, where the crystal structure of the RNA fragment can be deduced and computer programs are utilized to design novel binding compounds.

The compounds which can be tested and identified according to a method of the invention may be expression libraries, e.g., cDNA expression libraries, peptides, proteins, nucleic antibodies, small organic compounds, hormones, peptidomimetics, PNAs or the like (Milner, Nature Medicine 1 (1995), 879-880; Hupp, Cell 83 (1995), 237-245; Gibbs, Cell 79 (1994), 193-198 and references cited supra). Furthermore, genes encoding a putative regulator of a Kremen 1 and/or Kremen 2 polypeptide and/or which excert their effects up- or downstream a Kremen 1 and/or Kremen 2 polypeptide may be identified using, insertion mutagenesis using, for example, example, targeting vectors known in the art. Said compounds can also be functional derivatives or analogues of known inhibitors or activators. useful compounds can be for example Such transacting factors which bind to a Kremen 1 and/or Kremen 2 polypeptide or regulatory sequences of the gene encoding it. Identification of transacting factors can be carried out using standard methods in the art (see, e.g., Sambrook, supra). To determine whether a protein binds to the protein itself or regulatory sequences, standard native gel-shift analyses can be carried out. In order to identify a transacting factor which binds to the protein or regulatory sequence, the protein or regulatory sequence can be used as an affinity reagent in

standard protein purification methods, or as a probe for screening an expression library. The identification of nucleic acid molecules which encode polypeptides which interact with a Kremen 1 and/or Kremen 2 polypeptide described above can also be achieved, for example, as described in Scofield (Science 274 (1996), 2063-2065) by use of the so-called yeast "two-hybrid system". In this system the Kremen 1 or Kremen 2 polypeptide or a smaller part thereof is linked to the DNA-binding domain of the GAL4 transcription factor. A yeast strain expressing this fusion polypeptide and comprising a lacZ reporter gene driven by an the GAL4 recognized by promoter, which is appropriate transcription factor, is transformed with a library of cDNAs which will express plant proteins or peptides thereof fused to an activation domain. Thus, if a peptide encoded by one of the cDNAs is able to interact with the fusion peptide comprising a peptide of a Kremen 1 and/or Kremen 2 polypeptide, the complex is able to direct expression of the reporter gene. In this way the encoding Kremen 1 and Kremen nucleic acid molecules respectively, and the encoded peptide can be used to identify peptides and proteins interacting with a Kremen 1 and/or Kremen 2 polypeptide.

Once the transacting factor is identified, modulation of its binding to or regulation of expression of a Kremen 1 and/or Kremen 2 polypeptide can be pursued, beginning with, for example, screening for inhibitors against the binding of the transacting factor to a Kremen 1 or Kremen 2 polypeptide. Activation or repression of a Kremen 1 and/or Kremen 2 polypeptide could then be achieved in animals by applying the transacting factor (or its inhibitor) or the gene encoding it, e.g. in an expression vector. In addition, if the active form of the transacting factor is a dimer, dominant-negative mutants of the transacting factor could be made in order to inhibit its activity. Furthermore, upon identification of the transacting factor, further components in the signal cascade leading to activation (e.g. signal transduction) or repression of a gene

in the control of a Kremen 1 and/or Kremen involved be identified. Modulation of then can polypeptide activities of these components can then be pursued, in order to drugs and methods modulating develop additional for metabolism of protein degradation in animals. Thus, the present invention also relates to the use of the two-hybrid system as identification of activators the for above inhibitors of a Kremen 1 and/or Kremen 2 polypeptide.

The compounds isolated by the above methods also serve as lead compounds for the development of analog compounds. The analogs should have a stabilized electronic configuration and molecular conformation that allows key functional groups to be presented to a Kremen 1 and/or Kremen 2 polypeptide or its ligand in substantially the same way as the lead compound. In particular, the analog compounds have spatial electronic properties which are comparable to the binding region, but can be smaller molecules than the lead compound, frequently having a molecular weight below about 2 kD and preferably below about 1 Identification of analog compounds can be performed through use of techniques such as self-consistent field (SCF) interaction (CI) analysis, and normal configuration dynamics analysis. Computer programs for implementing these Computer-Assisted available; e.g., Rein, techniques are Modeling of Receptor-Ligand Interactions (Alan Liss, New York, 1989). Methods for the preparation of chemical derivatives and analogues are well known to those skilled in the art and are for example, Beilstein, Handbook of Organic described in, Chemistry, Springer edition New York Inc., 175 Fifth Avenue, New York, N.Y. 10010 U.S.A. and Organic Synthesis, Wiley, New York, USA. Furthermore, said derivatives and analogues can be tested for their effects according to methods known in the art; see also supra. Furthermore, peptidomimetics and/or computer aided design of appropriate derivatives and analogues can be used, for example, according to the methods described above.

Once the described compound has been identified and obtained, it is preferably provided in a therapeutically acceptable form.

invention also relates present the Accordingly, pharmaceutical composition comprising a nucleic acid molecule encoding a Kremen 1 and/or Kremen 2 polypeptide, a Kremen 1 and/or Kremen 2 polypeptide itself, recombinant vector (for antibody, activator/agonist, below), see examples, inhibitor/antagonist and/or binding partner of a Kremen 1 and/or Kremen 2 polypeptide and a pharmaceutically acceptable excipient, diluent or carrier.

Preferably, for therapeutic purposes, the Kremen 1 and/or Kremen 2 polypeptide is recombinantly produced by use of the nucleic acid sequences shown in Figures 1 and 2. Suitable vectors for recombinant expression are known to the person skilled in the art. Preferably, they are plasmids, cosmids, viruses, bacteriophages and other vectors usually used in the field of genetic engineering. Vectors suitable for use in the present invention include, but are not limited to the T7-based for expression in mammalian cells expression vector baculovirus-derived vectors for expression in insect cells. Preferably, the nucleic acid molecule of the invention the regulatory elements operatively linked the to invention that the guarantee recombinant vector of the transcription and synthesis of an mRNA in prokryotic and/or can be translated. The nucleotide that eukaryotic cells sequence to be transcribed can be operably linked to a promoter like a T7, metallothionein I or polyhedrin promoter. The host cells used for recombinant expression are prokaryotic eukaryotic cells, for example mammalian cells, bacterial cells, insect cells or yeast cells. The polypeptide is isolated from the cultivated cells and/or the culture medium. Isolation and purification of the recombinantly produced polypeptide may be including preparative means carried out by conventional chromatography and affinity and immunological separations using, e.g., an anti-Kremen 1 or 2 antibody, or, e.g., can be substantially purified by the one-step method described in Smith and Johnson, Gene 67; 31-40 (1988).

Examples of suitable pharmaceutical carriers etc. are well art and include phosphate buffered water, emulsions, such as oil/water various types of wetting agents, sterile solutions etc. Such carriers can be formulated by conventional methods and can be administered to the subject at a suitable dose. Administration of the suitable compositions may be effected by different ways, subcutaneous, intravenous, intraperetoneal, e.g. by intramuscular, topical or intradermal administration. The route of administration, of course, depends on the nature of the disease and the kind of compound contained The will pharmaceutical composition. dosage regimen be determined by the attending physician and other clinical factors. As is well known in the medical arts, dosages for any one patient depends on many factors, including the patient's size, body surface area, age, sex, the particular compound to be administered, time and route of administration, the kind and stage of the disease, e.g., tumor, general health and other. drugs being administered concurrently.

The delivery of the nucleic acid molecules encoding a Kremen 1 Kremen 2 polypeptide can be achieved by and/or application or, preferably, by using a recombinant expression vector such as a chimeric virus containing these compounds or a colloidal dispersion system. Direct application to the target site can be performed, e.g., by ballistic delivery, colloidal dispersion system or by catheter to a site in artery. The colloidal dispersion systems which can be used for delivery of the above nucleic acid molecules include macromolecule complexes, nanocapsules, microspheres, beads and lipid-based systems including oil-in-water emulsions (mixed), micelles, liposomes and lipoplexes, The preferred colloidal system is a

liposome. Organ-specific or cell-specific liposomes can be used in order to achieve delivery only to the desired tissue. The targeting of liposomes can be carried out by the person skilled in the art by applying commonly known methods. This targeting includes passive targeting (utilizing the natural tendency of the liposomes to distribute to cells of the RES in organs which contain sinusoidal capillaries) or active targeting (for example by coupling the liposome to a specific ligand, e.g., an antibody, a receptor, sugar, glycolipid, protein etc., by well known methods). In the present invention monoclonal antibodies are preferably used to target liposomes to specific tissues, e.g. tumor tissue, via specific cell-surface ligands.

Preferred recombinant vectors useful for gene therapy are viral vectors, e.g. adenovirus, herpes virus, vaccinia, preferably, an RNA virus such as a Retrovirus. Even more preferably, the retroviral vector is a derivative of a murine or avian retrovirus. Examples of such retroviral vectors which can be used in the present invention are: Moloney murine leukemia virus (MoMuLV), Harvey murine sarcoma virus (HaMuSV), murine mammary tumor virus (MuMTV) and Rous sarcoma virus (RSV). Most preferably, a non-human primate retroviral vector is employed, such as the gibbon ape leukemia virus (GaLV), providing a broader host range compared to murine vectors. Since recombinant retroviruses are defective, assistance is required in order to produce infectious particles. assistance can be provided, e.g., by using helper cell lines that contain plasmids encoding all of the structural genes of the retrovirus under the control of regulatory sequences within the LTR. Suitable helper cell lines are well known to those skilled in the art. Said vectors can additionally contain a gene encoding a selectable marker so that the transduced cells can be identified. Moreover, the retroviral vectors can be modified in such a way that they become target specific. This can be achieved, e.g., by inserting a polynucleotide encoding a sugar, a glycolipid, or a protein, preferably an antibody.

Those skilled in the art know additional methods for generating target specific vectors. Further suitable vectors and methods for in vitro- or in vivo-gene therapy are described in the literature and are known to the persons skilled in the art; see, e.g., WO 94/29469 or WO 97/00957.

In order to achieve expression only in the target organ, e.g., a tumor to be treated, the nucleic acid molecules encoding a Kremen 1 and/or Kremen 2 polypeptide can be linked to a tissue specific promoter and used for gene therapy. Such promoters are well known to those skilled in the art (see e.g. Zimmermann et al., (1994) Neuron 12, 11-24; Vidal et al.; (1990) EMBO J. 9, 833-840; Mayford et al., (1995), Cell 81, 891-904; Pinkert et al., (1987) Genes & Dev. 1, 268-76).

The present invention also relates to the use of the above the preparation compounds of invention for the composition treatment of for pharmaceutical associated with (a) aberrant expression of kremen 1, kremen 2 and/or genes involved into the Wnt signal cascade, and/or (b) aberrant activities or amounts of a Kremen 1, Kremen 2 and/or a involved into the Wnt signal cascade. polypeptide preferred embodiment, said disease is a tumor, preferably breast cancer, a colon carcinoma or a melanoma.

Finally, the present invention relates to the use of a nucleotide molecule encoding a polypeptide having a biological activity of Kremen 1 and/or Kremen 2, a Kremen 1 and/or Kremen 2 polypeptide, an activator/agonist of a Kremen 1 and/or Kremen 2 polypeptide or binding partner of said polypeptide(s) for the preparation of a pharmaceutical composition for inhibiting the Wnt signal cascade which might be useful for supporting regenerative processes in a patient, e.g. growth of tissue like muscle, hair, etc.

The following examples illustrate the invention.

Example 1

Isolation of cDNAs encoding Kremen 1 and 2, respectively

A mouse 13.5 day embryo cDNA library in the expression vector pCMV-SPORT2 (Gibco BRL) was used to prepare pools of about 250 colonies, and plasmid DNA from each pool was transiently transfected into 293T cells in 24-well plates using FuGENE 6 (Roche). After 48 hours cells were incubated with medium containing 1nM Dkk1-alkaline phosphatase (Dkk1-AP) fusion protein (Mao et al., Nature 411 (2001) 321-325) and processed for AP histochemistry. From 1500 pools, 2 positive pools were identified and single clones were isolated by sib selection. Sequencing analysis showed that they represent independent isolates of mkremen 2. A full length mouse kremen 1 clone was. library by PCR using published. isolated from the same nucleotide sequence data (Nakamura et al, Biochim. Biophys. Acta 1518 (2001), 63-72). The open reading frame of mkremen 1 and -2 was cloned into pCS2+ to generate pCS2-mkrm1 and -2. pCS-flag-mkrm2 was constructed by inserting a flag epitope after the signal peptide and was used as template to generate the pCS-flag-mkrm2\DeltaWSC by PCR.

Example 2

The binding of Kremen 1 and 2 to Dkk1 and Dkk2 shows high affinity and is physiologically relevant

For binding assays 293T cells were transfected (T) with mkrm1 or mkrm2 as indicated, incubated with recombinant Dkk1-alkaline phosphatase fusion protein (Dkk1-AP) or alkaline phosphatase (AP) and stained for bound AP activity. The results are shown in Figure 3.

As shown in Figure 4, luciferase Wnt reporter assays in 293T cells were done in 96 well plates at least in triplicates as described (Wu et al., Curr Biol 10 (2000), 1611-1614). Luciferase activity was normalized against Renilla activity using a commercial kit (Clonetech). Xdkk1= Xenopus dkk1 (Glinka, et al. Nature 391, (1998) 357-362); mkrm1,2= mouse kremen 1,2; wnt= mouse wnt1; fz= mouse frizzled8; lrp6= human lrp6 (Tamai, et al.. Nature 407 (2000) 530-535); Wnt luciferase reporter TOP-FLASH (Korinek et al. Science 275 (1997)1784-1787).

As shown in Figure 3, the binding of Dkk alkaline phosphatase fusion protein to Kremen 2 and Kremen 1, respectively, shows high affinity. Moreover, it could be shown that only Dkk1 and Dkk2 bind to Kremen but not Dkk3.

In an additional experiment, 293 kidney cells were transfected with the Wnt reporter (TOP-FLASH) with or without the genes indicated. Two days after transfection, the luciferase activity expressed was determined. As shown in Figure 4, cotransfection of Wnt and its receptor, frizzled (fz) results in stimulation of the Wnt signal cascade (see Figure 4, lane 1 versus lane 2) and cotransfection of dkkl and kremen 1 and kremen 2 leads to synergistic inhibition of this activation of the Wnt signal cascade. This effect is even more pronounced if wnt has been cotransfected with its receptor frizzled (fz) and the coreceptor lrp6. A very strong activation of the Wnt signal cascade (lane 8) can be observed. This activation can only inhibited by cotransfection with dkkl and kremen 1,2 (lanes 12 and 13) but not by transfection with the single genes (dkkl, lane 9; kremen 2, lane 10; kremen 1, lane 11).

Determination of the expression profile of kremen 1 and 2 in various tissues of mice

The expression of kremen 1 and 2 in various tissues of mice was studied by RT-PCR. RNA isolation from adult mouse organs and RT-PCR assays were carried out in the linear phase of amplification and with histone 4 primers as described (Glinka et al., Nature 389 (1997), 517-519) Other primers were: mkrm1 (f, GTGCTTCACAGCCAACGGTGCA; r, ACGTAGCACCAAGGGCTCACGT); mkrm2 (f, AGGGAAACTGGTCGGCTC; r, AAGGCACGGAGTAGGTTGC). Cycle no. were H4: 26 cycles; mkrm1: 35 cycles; mkrm2: 32 cycles. The results show that both kremens are expressed in all mouse tissues tested, but with varying expression level (Figure 5). Similar results were obtained using Xenopus embryos.

What is claimed is:

- 1. A diagnostic composition comprising
 - (a) at least one nucleic acid molecule which is capable of specifically hybridizing to the nucleotide sequence encoding Kremen 1 as depicted in Figure 1 and/or to the nucleotide sequence encoding Kremen 2 as depicted in Figure 2; or
 - (b) at least one ligand which is capable of specifically binding to a Kremen 1 and/or Kremen 2 polypeptide.
- 2. The diagnostic composition of claim 1, wherein the ligand is an antibody.
- 3. The diagnostic composition of claim 1, wherein the nucleic acid molecule has a length of at least 10 nucleotides.
- 4 The nucleic acid molecule or ligand of any one of claims 1 to 3, which is detectably labeled.
- 5. The nucleic acid molecule or ligand of claim 4, wherein the label is selected from the group consisting of a radioisotope, a bioluminescent compound, a chemiluminescent compound, a fluorescent compound, a metal chelate, or an enzyme.
- 6. The diagnostic composition of any one of claims 1 to 3, wherein the nucleic acid molecule or ligand are bound to a solid support.
- 7. Use of a nucleic acid molecule or ligand as defined in any one of claims 1 to 6 for the preparation of a diagnostic composition for the diagnosis of a disease associated with (a) aberrant expression of kremen 1 and/or kremen 2 and/or (b) aberrant activities or amounts of a Kremen 1 and/or Kremen 2 polypeptide.

- 8. Use according to claim 7, wherein the target to which the nucleic acid molecule hybridizes is an mRNA.
- 9. A method of diagnosing a disease associated with (a) aberrant expression of kremen 1 and/or kremen 2 and/or (b) aberrant activities or amounts of a Kremen 1 and/or Kremen 2 polypeptide in a subject comprising:
 - (a) determining (a) the amount of expression of kremen 1 and/or kremen 2 and/or (b) the amount of biologically active Kremen 1 and/or Kremen 2 polypeptide in a biological sample; and
 - (b) diagnosing a disease associated with (a) aberrant expression of kremen 1 and/or kremen 2 and/or (b) aberrant activities or amounts of a Kremen 1 and/or Kremen 2 polypeptide or a risk for the development of such disease based on an altered amount of expression of kremen 1 and/or kremen 2 and/or (b) an altered amount of biologically active Kremen 1 and/or Kremen 2 polypeptide compared to a control.
- 10 A method for identifying a binding partner to a Kremen 1 and/or Kremen 2 polypeptide comprising:
 - (a) contacting said polypeptide with a compound to be screened; and
 - (b) determining whether the compound effects an activity of said polypeptide or whether binding of the compound to said polypeptide has occured.
- activators/agonists identifying or. method for 11. A 2 inhibitors/antagonists of а Kremen 1 and/or Kremen polypeptide comprising the steps of:
 - (a) incubating a candidate compound with said polypeptide;
 - (b) assaying a biological activity, and
 - (c) determining if a biological activity of said polypeptide has been altered.

- 12. A method of identifying and obtaining a drug candidate for therapy of a disease associated with (a) aberrant expression of the gene encoding Kremen 1 and/or Kremen 2 and/or (b) aberrant activities or amounts of Kremen 1 and/or Kremen 2 comprising the steps of
 - (a) contacting a Kremen 1 and/or Kremen 2 polypeptide or a cell expressing said polypeptide, and optionally the corresponding ligand(s), in the presence of components capable of providing a detectable signal in response to binding to said drug candidate to be screened; and
 - (b) detecting presence or absence of a signal or increase of the signal generated, wherein the presence or increase of the signal is indicative for a putative drug.
- 13. An activator/agonist or inhibitor/antagonist of a Kremen 1 and/or Kremen 2 polypeptide or binding partner of said polypeptide(s) obtainable by the method of any one of claims 10 to 12.
- 14. A pharmaceutical composition comprising a compound which is capable of modulating the expression of a gene encoding Kremen 1 and/or Kremen 2 or the activity of Kremen 1 and/or Kremen 2 and a pharmaceutically acceptable excipient, diluent or carrier.
- 15. The pharmaceutical composition of claim 14, wherein the compound stimulates expression of the gene encoding Kremen 1 and/or Kremen 2 or the activity of Kremen 1 and/or Kremen 2.
- 16. The pharmaceutical composition of claim 15, wherein the compound is a nucleotide molecule encoding a polypeptide having a biological activity of Kremen 1 and/or Kremen 2, a Kremen 1 and/or Kremen 2 polypeptide, an activator/agonist or inhibitor/antagonist of a Kremen 1 and/or Kremen 2 polypeptide or binding partner of said polypeptide(s) obtainable by the method of any one of claims 10 to 12.

- 17. Use of a compound as defined in claim 16 for the preparation of a pharmaceutical composition for the treatment of a disease associated with (a) aberrant expression of kremen 1, kremen 2 and/or a gene involved into the wnt signal cascade and/or (b) aberrant activities or amounts of a Kremen 1, Kremen 2 and/or polypeptide involved into the Wnt signal cascade.
- 18. Use according to claim 7 or 17, wherein the disease is a tumor or a disease of the kidneys, bones and eyes.
- 19. Use of a nucleotide molecule encoding a polypeptide having a biological activity of Kremen 1 and/or Kremen 2, a Kremen 1 and/or Kremen 2 polypeptide, an activator/agonist of a Kremen 1 and/or Kremen 2 polypeptide or binding partner of said polypeptide(s) for the preparation of a pharmaceutical composition for inhibiting the Wnt signal cascade.
- 20. Use according to claim 19 for supporting regenerative processes.

Abstract

Compositions for diagnosis and therapy of diseases associated with aberrant expression of kremen and/or wnt

The present invention relates to a composition useful for the diagnosis of diseases associated with aberrant expression of the gene encoding the receptor Kremen 1 and/or Kremen 2 e.g. tumors or diseases of the kidneys, bones and eyes. The present invention also relates to a pharmaceutical composition containing a compound which is capable of modifying (a) the expression of the gene encoding Kremen 1 and/or Kremen 2 or (b) the activity of the Kremen 1 and/or Kremen 2 receptor.

mkrml hkrml mkrm2 hkrm2	ATGGGGACAC ATGGGGACAC	T G G C G C G C C T G C A A A G C C C T G C A	CGGCTTCCTC	CTCGCGCT CTCGC;CCT CTCCTCT	29 29 37 40
mkrml hkrml mkrm2 hkrm2	GCTCTCGGCC GCTCTCGGCC TCCCATTGCT TCCTCCGCT	GCTGCGCTCA GCGGCTGA GCTGCGGCTG GCTGCAGCCG	CTCTGG CGCTGG CACGGGGCCT CGTGGGGCCT	C G G C C G G C C G G C C C G G C C G	65 65 77 80
mkrml hkrml mkrm2 hkrm2	C C C C C C C C C C C C C C C C C C C	CCCCCCTCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCC	GCCCCGAGTG GCGCCGAGTG TCCGAATG	CTTCACAGGG TTTCACAGGTG CTTCCAGGTG	105 ⁻ 105 114 117
mkrm1 hkrm1 mkrm2 hkrm2	AACGGTGCAG AATGGTGCGG AACGGCGCTG AATGGGGCTG	ATTACAGGGGATTATAGGGGACTACCGGGGAGGACTACCGGGG	AACACAGAGC AACACAGAAC CGACCAGAAC CGACCAGAAC	T G G A C A G C G C T G G A C A G C C C C G C A C C G G C C	145 145 154 157
mkrml hkrml mkrm2 hkrm2	TGCA, AGG TACA.AGG CACGCGGAGG CGCGCGGGGC	T G G G A A G C C A, C G G G A A G C C A T G G A C G C C C T G G G C C G C C G	TGTCTGTTCT TGTCTGTTTT TGTCTTTTCT	G G A A C G A G A C G G A A C G A G A C G G G A C C A G A C G G G A C O A G A C	182 182 194 197
mkrm1 hkrm1 mkrm2 hkrm2	TTTCCAGCAT TTTCCAGCAT ACAGCAGCAC GCAGCAACAC	CCGTACAACA CCATACAACA AGCTACAGCA AGCTACAGCA	CGCTGAAGTA CTCTGAAATA GCGCCAGCGA GCGCCAGCGA	C C C C A A C G G G C C C C A A C G G G C C C C	222 222 234 237
mkrml hkrml mkrm2 hkrm2	G A A G G A G G A G G A G G G G G C C C G C T G G G G G T C G C T G G G G C	TGGGGGAGCA TGGGTGAGCA TGGGTGCGCA TGGGCGGCA	CAATTATTGC CAACTATTGC TAACTTCTGT CAACTTCTGC	AGAATCCAG AGAAATCCAG AGGAACCCAG GGTAACCCAG	262 262 274 277
nkrm1 nkrm1 nkrm1 hkrm1	ATGGAGACGT ATGGAGACGT ACGGTGATGT ACGGTGACGT	GAGCCCTTGG GAGCCCCTGG GCAGCCCTGG GCAGCCGTGG	TGCTACGTGGTGCTGCTATGTGGTGCTACGTGGTGGTGGTGGTGGTGG	CCGAGCATGA CAGAGCACGA CAGAGACAGA CTGAGACAGA	302 302 314 317
mkrml hkrml mkrm2 hkrm2	GGACGGAGTC GGATGGTGTC AGAGGGCATC GGAGGGCATC	T A C T G G A A G T T A C T G G A A G T T A C T G G C G C T T A C T G G C G C T	ACTGTGAAAT ACTGTGAGAT ACTGTGATAT ACTGCGACAT	TCCTGCTTGCACCTGCCCCACATGT	342 342 354 357
mkrml hkrml mkrm2 hkrm2	CAGATGCCTG CAGATGCCTG CACATGCCTG CACATGCCAG	G A A A C C T T G G G A A A C C T T G G G G T A C C T G G G G C T A C C T G G G	CTGCTACAAG CTGCTACAAG GTGCTTCGTG ATGCTTTGTG	GATCATGGAA GATCATGGAA GACTCTGGG GACTCAGGG	382 382 394 397
mkrm1 hkrm1 mkrm2 hkrm2	ACCCACCTCC ACCCACCTCC CACCCCCTGC CACCCCCAGC	TCTCACGGCC TCTAACTGGC TCTCAGTGGT CCTCAGCGGC	ACCAGTAAAA ACCAGTAAAA CCCAGTGGCA CCCAGCGGCA	CCTCTAACAA CGTCCAACAA CCTCCACAAA	
mkrm1 hkrm1 mkrm2 hkrm2	GCTCACCATA ACTCACCATA GCTCACTGTC GCTCACGGTC	CAAACCTGTA CAAACTTGCA CAAGTGTGCC CAGGTGTGCC	TCAGCTTCTG TCAGTTTTTG TTCGATTGTG TACGCTTCTG	TCGGAGTCAG TCGGAGTGAAG CCGCATGAAG	462 462 474 477
mkrml hkrml mkrm2 hkrm2	AGATTCAAGT AGGTTCAAGT GGCTACCAGC GGCTACCAGC	1:96666666		TATGCCTGCTTATGCTTATGCTTATGCCTTACCCT	517
mkrml hkrml mkrm2 hkrm2	TCTGTGGGAA TCTGTGGAAA TCTGTGGCTC	CAATCCTGAC CAATCCTGAT TGAAAGTGAC TGAAAGCGAC	TA C'T G G A A G C TA C T G G A A G T C T G G C C C G C G C T G G C C C G G G	G A C G C C T C G C	542
mkrml hkrml mkrm2 hkrm2	G G C C A G C A C C A G C C A G T A C C C C C T G C C A C C C C C C C C A C C	GAATGCAACA	GCGTCTGCTT	CGGGGACCAC GGGGGATCAC TGGCCACCCA	582 582 594 597

Multiple-alignment of mouse and human kremen DNAs (3-1)

			Y		
mkrm1	ACCCAGCCT	G C G G G G A	CGGCAGATT	A TCCTCTTTG	622
hkrm1	ACCCAACCCT	G T G G G G A	TGGCAGATC	ATCCTCTTTTG	622
mkrm2	GGCCAGCTCT	G T G G A G G G A	TGGACGACTA	GGCATCTATG	634
hkrm2	GGCCAGCTGT	G T G G C G G G A	TGGGCGCTG	GGCGTCTATG	637
mkrm1 hkrm1 mkrm2 hkrm2	ACACITCTCGT ACACTCTCGT AAGTGTCTGT AAGTGTCGGT	G G G C C T G C G G G C T C C T G C G G G C T C C T G C	G G T G G G A A C T G G T G G G A A C T C A G G G A A A C T C A G G G G A A C T	ACTCAGCGAT ACTCAGCCAT GGTCGGCTCC GGACAGCGCC	662 662 674 677
mkrm1	GG GA G C C G T G	GTGTACTCCC	CTGAGTTCCC	T G A C A C C T A C	702
hkrm1	GT C T T C T G T G	GTGTATTCCC	CTGAGTTCCC	G G A C A C C T A T	702
mkrm2	T C A A G G A G T C	ATCTACTCCC	CGGATTTCCC	G G A T G A G T A T	714
hkrm2	T C A G G G C G T C	ATCTACTCCC	CGGATTTCCC	G G A C G A G T A C	717
mkrm1	GCCACTGGGA	GAGTCTGCTA	CTGGACCATC	C G G G T T C C A G	742
hkrm1	GCCACGGGGA	GGGTCTGCTA	CTGGACCATC	C G G G T T C C G G	742
mkrm2	GGA,GCAGACC	GGAACTGCAG	CTGGGTATTG	G G C C A A C T G G	754
hkrm2	GGGCCGGACC	GGAACTGCAG	CTGGGCCCTG	G G C C C G C C A G	757
mkrm1 hkrm1 mkrm2 hkrm2	GAGCCTCTCGA GGGCCTGTGC. GCGCTGTGC.	CATCCATTTC CATCCACTTC . TAGAAGTC . TGGAGCTC	AACTTCACCCAGCCAGCTTCCGCCAGCTTCCGCC	T G T T T G A T A T T A T T T G A C A T T C T T C G A G T T T C T T C G A G C T	782 782 791 794
mkrm1 hkrm2 hkrm2	CAGGAGTCT CAGGGACTCG GGCMGATTCT GGCCGACCC	G C A G A C A T G G G C G G A C A T G G G G A G A C C G G C C G G G A C C G G G	T G G A G C T G C T T G G A G C T T C T T G G A G C T A C G T G G A G C T G C G	GGACGGCTAC GGATGGCTAC CGACGTCT	822 822 829 832
mkrm1	ACCCACCGCG	TCCTGGTCCG TCCTAGGCCG ACCTACTCCG GCCTGCTCCG	G C T C A G T G G G	A G G A G C C,G C C	862
hkrm1	ACCCACCGTG		C T T C C A G G G G	A G G A G C C G C C	862
mkrm2	. CGTCCGGCA		T G C C T T C G A C	G C C C C A T C	868
hkrm2	CTTCGGGCA		C G G C T T C G A T	G G C G C C G C C.	871
mkrm1	C G C C T C T G T C C C C C C C C C C C	TTTCAATGTC	T C T C T G G A T T	TTGTCATTT	902
hkrm1		CTTCAACGTC	T C T C T G G A C T	TCGTCATCTT	902.
mkrm2		GGGACCGGTG	C G C C T G C G C A	CTGCTGCGCT	908
hkrm2		CGGGCCGGTG	C G C C T G G C A	CTGCCGCT	911
mkrm1	GTATTTCTTC	T C T G A T C G C A T C T G A T C G C A T T C C G C A G C G T T C C G A A G C G	TCAATCAGGC	CCAGGGATTT	942
hkrm1	GTATTTCTTC		TCAATCAGGC	CCAGGGATTT	942
mkrm2	GCTGCTCACC		ACGCAAGAGG	CCATGCTC	946
hkrm2	GCTGCTCACC		ACGCGCGCGG	CCACGCGC	949
mkrml	GCTGTGTTGT	A C C A A G C A C	CAAGGAGAA	C.C.C.C.A.C.A.G.G.C.C.A.C.A.G.G.A.G.G.A.C.A.G.T.A.C.A.G.T.A.G.G.A.G.C.G.C.G.C.G.C.G.C.G.C.G.C.G.C	982
hkrm1	GCTGTTTTAT	A C C A A G C C G T	CAAGGAAGAA		982
mkrm2	AAGGCTTTCGC	G C T C A C C T A C	CGCGGGCTGC		986
hkrm2	AAGGCTTCGC	G C T C A C C T A C	CGCGGGCTGC		989
mkrml	AGAGAGCTGC	TGTCAACCAG	ACCCT.GGCAG	A G G.T G A T C A C	1022
hkrml	AGAGGCCGG	TGTCAACCAG	ACGCT.GGCGA	A G G T G A T C A C	1022
mkrm2	GGAGGAGA	GCATCTCCAG	AGGATT.CAA	C T G A G A G T C T	1025
hkrm2	TGAGGACCA	GAGGCCCCCG	AGGGCT.C	C C C A G A C C C	1028
mkrml	0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	AACCTCAGTG	T C A G C G C T G C	CCACTCCTCC	1062
hkrml		AACCTCAGTG	T C A G C G C T G C	CCGGTCCTCC	1062
mkrm2		CCCGATGGG	C T A A C G C G A G	CTGCAGCCCC	1065
hkrm2		CTCGACGG	C C A A C G T G A G	CTGCAGCCCC	1068
mkrm1	AAAGTCCTCT	ATGTCATCAC	C C C C A G C C C C	A G C C A C C C T C	1102
hkrm1	AAAGTCCTCT	ATGTCATCAC	C A C C A G C C C C	A G C C A C C C A C	1102
mkrm2	AAGCCC	GGAGGTGCAC	A G C C T T C G A T	A G G T G C C G A	1101
hkrm2	AGGCCT	GGGGGTCC	C G C C C C A T	T G G G C C C G G	1104
mkrm1	CGCAGACTGC	CCCAGGTAGC	C A T T C C T G G G G A A T T C C T G C G G C T G C T G T G T G T	CACCATCAGT	1142
hkrm1	CTCAGACTGT	CCCAGGTAGC		CCCCAT	1142
mkrm2	GTCTTCTCGA	CCGTGACGC		CTGCTGCTGT	1141
hkrm2	GTCTTCTCGA	CGGTGACGGC		CTGCTGCTGC	1144
mkrm1	TGGGGCCAAC	AGCCACAGAG	T G G A A G G A T G	GACTGTGTAC	1182
hkrm1	GGGGGCTGGA	AGCCACAGAG	T T G A A G G A T G	GACAGTCTAT	1182
mkrm2	TGCTCCTGTC	CCTACTGCGT	T T G C T G C G T C	GACGG	1176
hkrm2	TGCTGCTGGG	GCTGCTGCGT	C G G C T G C G C C	GACGGTGCGG	1184

Multiple-alignment of mouse and human kremen DNAs (3-2)

```
CCCTCCTGAT CCTCACAGTC ACAGCAGTTG
CTCTCTCATA CCTCACAGTC ACAGCCATTG
CAGGGCCTGA GGGCGGA.CC GGTGGAGCTG
                  GGCCTGGCGA
GGTCTGGCAA
mkrml
                                                                                                                                                                                                 1222
hkrml
                                                                                                                                                                                                 11,81
mkrm2
                   GGCGCTGGGG
hkrm2
                                                                                                                                                    AAT CTCATCG
AATCCCATCG
CATGGGACCT
GCTGGGGGCT
                                                              T C T T C T G C A T

 C T T C T G G A C

C C A G G A A A A G

C C G G G A A A A G
                                                                                                         GT CA CATTTA
GT CA CATTCA
GGT CTCTGGC
GGCCCCGGC
                  T C G G A A A G A T
T A G C A A A G A T
T C T G C T G G C T
T C T G C T G G C T
                                                                                                                                                                                                 1262
mkrml
                                                                                                                                                                                                 1262
hkr.ml
                                                                                                                                                                                                 1221
 mkr:m2
hkrm2
                                                                                                                                                    T C G T C A G C C T
T C A T C A A C C A
T G G T A C C G C C
T G G T A C C A A C
                                                                                                          TTAGGGACTG
TTAGGGATTG
CTGGGCTGTG
CTGGGCTGTG
                                                              T CA G G A G A C C
T CA G G G G A G C
G C G G G A G A A G
C CA G G A G A A G
                  A G T C C C T G C A
T G T T C C T G C T
T C E C G G G G C C
T C C A G G G C C
                                                                                                                                                                                                 1302
1302
hkrm1
                                                                                                                                                                                                 1261
mkrm2
                                                                                                                                                                                                 1303
hkrm2
                                                                                                                                                    TATGAAGCTTT
TACAAGCCTTC
CAGGGGACTC
CCGGGGGACCC
                                                             GAGATATCTG
GGGAAATCTG
GGTGGCCCTG
GGTGGCCTTG
                   O G G G C T T C T G
G G G A C T T C G G
G G G G C C G A G G
A G C C C G A G G
                                                                                                          GACGATTTTC
                                                                                                                                                                                                 1342
 mkr.m1
                                                                                                          GAGCATTTTT
CCCTGTCCCC
CCTGCTCCC
                                                                                                                                                                                                 1342
1301
 hkrm1
 mkrm2
 hkrm2
                                                                                                          TTAAGAAGAA
TTAAGAAGAA
CGGGCTACCG
                                                                                                                                                     G C T C A A G G G T
A C T C A A G G G T
T C C C C T G A G T
G C C T C T G A G T
                                                              A T C T C C A T C T
A T T T C C A T C T
G G T C C T G C T G
G G T T C T G C C G
                                                                                                                                                                                                  1380
                   C C A . . . C T A C A C C A . . . C T IT C A T C A G G C T G A G C C A G G C T G A G
 mkrml
                                                                                                                                                                                                 1380
 hkrml
                                                                                                                                                                                                  1341
 mkrm2
 hkrm2
                                                                                                                                                     GTGAGTGACT
GTGAGTGACT
GTCTCTGCTC
                                                                                                          CAATCCCCTC
CAATCCCCTT
GCGCTCGCTC
                   CAGAGTCAAC
CAGAGTCAAC
GCCTCCAGG
GCCTCCAGG
                                                            A A G A T G A C C G
A A G A T G A C C G
A G A G C T C C T T
A G A G C T C C C T
                                                                                                                                                                                                  1420
 mkrm1
                                                                                                                                                                                                 1420
1381
 hkrml
 mkrm2
 hkrm2
 mkrml
                    GA - - -
                                           1422
                   AA---
TCTGA
TCTGA
                                           1422
1386
1428
 hkr.ml
 mkrm2
 hkrm2
```

Multiple-alignment of mouse and human kremen DNAs (3-3)

```
1 MAP PAARLALLSAAALTLAARPAPEPRS.GP...ECFTANGADYRGTQSWTALQG
1 MAP PAARLALLSAAALTLAARPAPEPGL.GP...ECFTANGADYRGTQNWTALQG
1 MGTPHLQGFLLLFPLLLR.LEGASAGSLESPGLSECFQVNGADYRGEQNETGPRG
1 MGTQALQGFLFLLFLPLLQPRGASAGSLESPGLSECFQVNGADYRGEQNRTGPRG
mkrm1
hkrm1
mkrm2
 hkrm2
                        52 .GEPCLFWNETFQHPYNELKYPNGEGGLGEHNECRNPDGDVSPWCYVAEHEDGWY
52 .GEPCLFWNETFQHPYNELKYPNGEGGLGEHNECRNPDGDVSPWCYVAEHEDGWY
55 AGEPCLFWDQTQQHEYSSASDPQGRWGLGAHNECRNPDGDVQPWCYVAETEEGEY
56 AGEPCLFWDQTQQHSYSSASDPEGRWGLGAHNECRNPDGDVQPWCYVAETEEGEY
mkrm1
hkrm1
mkrm2
mkrml 106 WWYCEIPACOMPGNLGCYKDHGNPPPLEGTSKTSNKLTEOTCESFCRSOREKFAG.
hkrml 106 WWYCEIPACOMPGNLGCYKDHGNPPPLEGTSKTSNKLTEOTCESFCRSOREKFAG.
mkrm2 110 WWYCEIPACOMPGNLGCYKDHGNPPPLEGTSKTSNKLTEOTCESFCRSOREKFAG.
hkrm2 111 WWYCEIPECEMPGYLGCEVDSGAPPALEGPSGTSTKLTEOVCERFCRMKGWQLAG.
mkiml 161 MESGYACFCGNNPDYWMHGBAASTECNSWCFGDHTQPCGGDGRALLEDTLVGACG
hkiml 161 MESGYACFCGNNPDYWMYGBAASTECNSWCFGDHTQPCGGDGRALLEDTLVGACG
mkim2 165 WEAGYACFCGSESDLAMGREAPATDCDQECFGHPGQLCGGDGRAGAWEVSVGSCQ
hkim2 166 WEAGYACFCGSESDLAMGRAPATDCDQECFGHPGQLCGGDGRAGAWEVSVGSCQ
mkrml 216 GNYSAMANVÄYSPDFPDTYNTGRVCYWTORVPGASKEHFNFTLFDORDSADMÖEL
hkrml 216 GNYSAMSSVVYSPDFPDTYNTGRVCYWTORVPGASKOHFNFTLFDORDSADMÖEL
mkrm2 220 GNWSAROGVTYSPDFPDEYGPDRNCSWVGGOLGAV.DELTFRLFERADSRDROEL
hkrm2 221 GNWBAROGVEYSPDFPDEYGPDRNCSWANGPPGAN.DELTFRLFERADPRDROEL
mkrml 271 LDGYTHRULVRLSGRSRPP.LSFNWSLDFWHLYFFSDRINQAQGFAVLYQBTKEE
hkrml 271 LDGYTHRULARDHGRSRPP.LSFNWSLDFWHLYFFSDRINQAQGFAVLYQBWKED
mkrm2 274 RDVSSGNELRAFDGAHPPPPGPLRURTAADELTFRSDARGHAQGFAHTYRGLQDT
hkrm2 275 RDBASGSELRAFDGARPPPSGPLRUGTAADELTFRSDARGHAQGFAHTYRGLQDA
mkrm1 325 PPQERPAVNQTPAEVITEQANLSVSAAHSSKVLYVEEPSPSHPPQTAPGSHSWAP
hkrm1 325 PPQERPAVNQTVAEVITEQANLSVSAARSSKVLYVIEPSPSHPPQTVPGSNSWAP
mkrm2 329 WE...GRASPEDSTESLAGDPDGAN...ASCSP....XPG...AA
hkrm2 330 AE....PPEAPEGSAQTPAAPEDGAN...WSCSP....XEG...AP
mkrm1 427 .DLRDCROPGASGDIWTER YEPSTTESEFKKKLKGOSO..QDDRNPLVSD-----
bkrm1 427 .DLRDCROPGISGEIWSETS YEESTSEFKKKLKGOSO..QDDRNPLVSD-----
mkrm2 403 LAMGPSRGPGRS...WAVWYDRPRGVAEPCPGDSQAEGPAAGYRPLSASSQSSL
hkrm2 417 PALGASEGPRS...WAVWYQQPRGVAEPCSPGDPQAEGSAAGYRPLSASSQSSL
 mkrm1 474 -----
 hkrm1 474 -----
mkrm2 455 RSLVSAL
hkrm2 469 RSLESAL
```

Multiple-alignment of mouse and human Kremen proteins

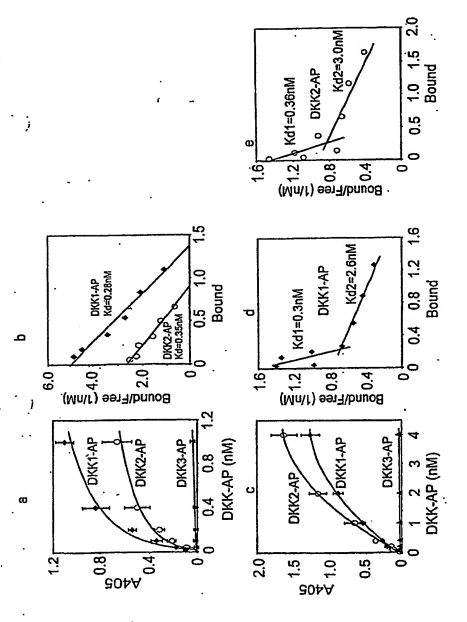


Fig.

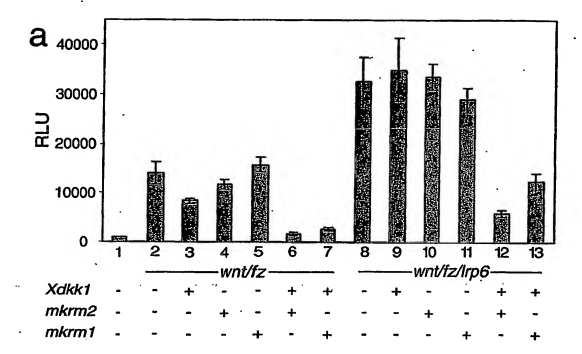
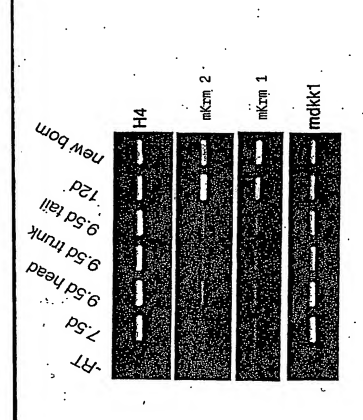
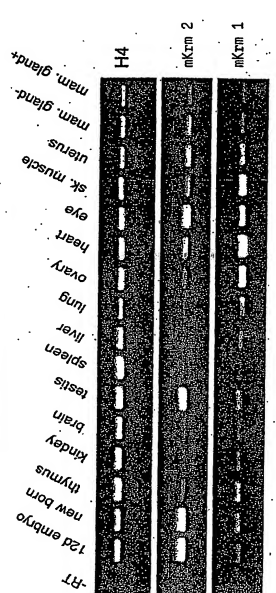


Fig. 4





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